Efficacy of ethyl-eicosapentaenoic acid in bipolar depression: randomised double-blind placebo-controlled study*

SOPHIA FRANGOU, MICHAEL LEWIS and PAUL MCCRONE

Background  Epidemiological and clinical studies suggest that increased intake of eicosapentaenoic acid (EPA) alleviates unipolar depression.

Aims  To examine the efficacy of EPA in treating depression in bipolar disorder.

Method  In a 12-week, double-blind study individuals with bipolar depression were randomly assigned to adjunctive treatment with placebo (n = 26) or with 1 g/day (n = 24) or 2 g/day (n = 25) of ethyl-EPA. Primary efficacy was assessed by the Hamilton Rating Scale for Depression (HRSD), with changes in the Young Mania Rating Scale and Clinical Global Impression Scale (CGI) as secondary outcome measures.

Results  There was no apparent benefit of 2 g over 1 g ethyl-EPA daily. Significant improvement was noted with ethyl-EPA treatment compared with placebo in the HRSD (P = 0.04) and the CGI (P = 0.004) scores. Both doses were well tolerated.

Conclusions  Adjunctive ethyl-EPA is an effective and well-tolerated intervention in bipolar depression.

Declaration of interest  The study (excluding attendance or presentations at international conferences) was supported by Laxdale Ltd, supplier of the ethyl-EPA preparation used in it.

In spite of the often dramatic nature of mania, the depressive phases of bipolar disorder can contribute most to poor outcome (MacQueen et al, 2001). Treatment is both understudied and clinically complicated (Compton & Nemeroff, 2000). Interest has grown in the potential role of omega-3 fatty acids such as eicosapentaenoic acid (EPA), which are found in certain plants and marine animals such as ‘oily’ fish. A possible role in the treatment of bipolar depression is suggested by studies of fish consumption (Hibbeln, 1998; Noaghiul & Hibbeln, 2003), blood fatty acid biochemistry (Adams et al, 1996) and clinical trials (Horrobin & Peet, 2001; Nemets et al, 2002; Puri et al, 2002). There is some evidence that they might prolong inter-episode remission in people with bipolar disorder (Stoll et al, 1999). Our aim therefore was to examine the efficacy and tolerability of ethyl-EPA as an adjunctive treatment for bipolar depression.

METHOD

Study design  The study was a single-centre, 12-week, double-blind randomised comparison of ethyl-EPA at 1 g or 2 g/day vs. placebo (paraffin oil) as adjunctive treatment in out-patients with bipolar depression. The decision to examine the efficacy of two doses of ethyl-EPA was based on previous studies that had found 2 g/day of ethyl-EPA to be the optimal dose for schizophrenia (Peet et al, 2002) and 1 g/day for unipolar depression (Horrobin & Peet, 2001).

Because of lack of data on the efficacy of ethyl-EPA in bipolar depression at the time of initiation of the study, formal sample size calculations were not possible. This study was therefore not powered to detect changes between the three treatment groups but to allow preliminary data to be collected regarding treatment effect size (if any) for planning future studies. The study was conducted at the Institute of Psychiatry, London, according to the principles of the Declaration of Helsinki and was approved by the local ethics committee. Participants were recruited following referral from their treating physicians or through advertisements in patient groups’ newsletters. After a complete description of the study, written informed consent was obtained from all participants and signed agreement was obtained from their treating physicians.

Participants were then screened to confirm their eligibility. Eligible participants were males or females between the ages 18 and 70 years who met criteria for bipolar disorder I or II as set out in the DSM–IV (American Psychiatric Association, 1994) and as determined by personal interview using the research version of the Structured Clinical Interview for DSM–IV (First et al, 1994).

Participants were also required to score at least 10 on the 17-item Hamilton Rating Scale for Depression (HRSD–17 Hamilton, 1960). Individuals were not included if: there was evidence of alcohol or illicit substance dependence, as defined by DSM–IV criteria, over the preceding 6 months; the severity of their bipolar disorder was such that participation in a clinical trial was not appropriate because of risk of imminent suicide or admission to hospital; there was a history of poor adherence to treatment and poor attendance at appointments; there was a concurrent medical condition or medication that could have accounted for the depressive episode; they had clinically significant abnormalities on routine biochemistry and haematology tests; they were on anticoagulants; they had known allergies to the ingredients of the study medication; they had taken fatty acid supplements or had been exposed to study medication in the preceding 12 weeks; or, in the case of women, they were pregnant or lactating, or of child-bearing potential and not taking adequate contraceptive precautions.

Eligible participants underwent a baseline assessment using the HRSD, the Young Mania Rating Scale (YMRS; Young et al, 1978) and the Clinical Global Impression Scale (CGI; Guy, 2000). Information about their concomitant medication was also recorded at baseline. There were no restrictions to the type and dose of psychotropic medication that they were receiving upon study entry. Participants were randomised only if existing psychotropic medication...
outcomes of the ethyl-EPA and placebo groups we used linear regression analysis on an intent-to-treat basis. With the regression models we were able to control for baseline scores in a similar way to using analysis of variance but with the added benefit of being able to use bootstrapping techniques to generate robust confidence intervals in the presence of data that followed a non-normal distribution. Bootstrapping involves resampling from the original data a sufficient number of times (5000 in this study) in order to approximate the population from which the sample is drawn; this does not involve prior assumptions as to the form of this distribution. In the results that follow the mean difference and standard errors (s.e.) are reported along with the bootstrapped 95% confidence intervals of the difference.

The Cohen’s $d$ effect sizes were also calculated to determine the magnitude of the differences between the treatment and placebo groups in depression and mania ratings (Cohen, 1988).

The study was funded by Laxdale Ltd, who collaborated with the authors on study design but were not involved in data collection, analysis or interpretation, writing the report or in the decision to submit for publication. Study materials were packaged and masked by the Clinical Trial Supplies Company and adverse events were monitored by Clintrials Research Ltd; neither was involved in any other aspect of the study.

RESULTS

A total of 93 people were screened for eligibility. The flow of potential participants is shown in Fig. 1. Of these, 18 were ineligible because of an incorrect diagnosis ($n=3$), an HRSD score below 10 ($n=4$), concurrent substance misuse ($n=1$), medical conditions ($n=2$), frequent medication changes ($n=5$) and withdrawal of consent prior to randomisation ($n=3$). The remaining 75 people were enrolled in the study between January and December 2001 and follow-up was completed at the end of March 2002. The clinical and demographic characteristics of the study participants are shown in Table 1. Participants were well matched in terms of their clinical and demographic characteristics. Table 2 summarises participants’ medication at study entry.

In total, nine individuals stopped taking the study medication, six from the placebo group and three of those randomised to receive 2 g/day ethyl-EPA. For all but two of these lack of efficacy was the reason for discontinuing study medication. Of the other two, one misunderstood the study protocol and stopped study medication when their concomitant medication was changed and the other did not like the appearance of the study medication. However, only four individuals (two in the placebo and two in the 2 g/day ethyl-EPA groups) failed to complete their assessments at study end-point; for these four the HRSD, YMRS and CGI scores were extrapolated using the last-observation-carried-forward method. Results were analysed on an intent-to-treat basis, including participants who stopped the study medication.

Table 3 summarises the mean and standard deviations of the participants’ scores at study entry and end-point. Figures 2 and 3 show the changes in HRSD and YMRS scores across groups between baseline and study end-point. There were no group differences in episode duration at the time of study entry ($F=3.9$, $d.f.=2$, $P=0.6$) or in the baseline scores on the HRSD ($F=0.8$, $d.f.=2$, $P=0.4$), YMRS ($F=0.6$, $d.f.=2$, $P=0.5$) or CGI ($F=0.5$, $d.f.=2$, $P=0.5$).

Exploration of initial data revealed no difference between the two ethyl-EPA groups in terms of end-point HRSD, YMRS and CGI scores. Data analysis was performed with the two active treatment groups combined.

(a) In terms of the main outcome measure, the mean HRSD score at the week 12 visit was 3.3 (s.e.=1.40) points lower for the ethyl-EPA groups (bootstrapped 95% CI $-6.1$ to $-0.2$, $P=0.03$). The overall HRSD effect size calculated from the difference between baseline and end-point measurements was 0.34 by Cohen’s $d$.

(b) The mean YMRS score at the week 12 visit was 3.3 (s.e.=2.2) points lower for the ethyl-EPA group compared with the placebo group (bootstrapped 95% CI $-8.6$ to $1.6$, $P=0.17$). The
During the trial, 26 of the 75 randomly assigned participants were included in the placebo group, 21% of the 1 g/day ethyl-EPA group and two in the 2 g/day ethyl-EPA group. There was no difference between the groups in these two types of side-effects ($\chi^2 = 1.2$, d.f. = 2, $P = 0.59$). There were also reports of isolated side-effects: two people in the placebo group reported constipation, there was one report of nausea and one of flatulence in the 1 g/day ethyl-EPA group and one report of an unpleasant taste in the 2 g/day ethyl-EPA group.

At study end-point the 71 participants (95% of the randomised sample) who completed their assessments were asked whether they thought they had received active treatment or not. There were no group differences regarding participants’ ability to guess their group allocation ($\chi^2 = 1.2$, d.f. = 2, $P = 0.5$); only 23% of the placebo group, 21% of the 1 g/day ethyl-EPA and 24% of the 2 g/day ethyl-EPA groups guessed their allocation correctly.

**DISCUSSION**

**Efficacy of ethyl-EPA in bipolar depression**

Treatment of bipolar depression with adjunctive ethyl-EPA resulted in improved clinical outcomes compared with placebo in terms of reduction in HRSD and CGI scores. Improvement was not significantly different in participants treated with 2 g/day as opposed to 1 g/day of ethyl-EPA.

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**Fig. 1** CONSORT diagram showing the flow of participants through each stage of the trial.

**Fig. 2** Hamilton Rating Scale for Depression (HRSD) scores in the placebo (n = 26) and combined ethyl-eicosapentaenoic acid (EPA) groups (n = 49) at baseline ( ), week 4 ( ) and week 12 ( ). The thick black line represents the mean, the whiskers are the standard deviations and the box is the range.

**Fig. 3** Young Mania Rating Scale (YMRS) scores in the placebo (n = 26) and combined ethyl-eicosapentaenoic acid (EPA) groups (n = 49) at baseline ( ), week 4 ( ) and week 12 ( ). The thick black line represents the mean, the whiskers are the standard deviations and the box is the range.
Baseline and end-point ratings on the YMRS were not significantly different among the three groups. Although there have been reports of hypomania during treatment with a different preparation of omega-3 fatty acids (Kinrys, 2000), we found no evidence that treatment with ethyl-EPA precipitates polarity changes in people with bipolar disorder.

**Methodological considerations**

There are several methodological issues that are worth considering. The placebo response rate in clinical trials of bipolar depression is high, with a pooled average of 29% (Keck et al., 2000). To control for the influence of psychosocial factors, we kept the number of assessments and contacts with the research team at a minimum to minimise the possibility that benefits from treatment could result from increased contact with health professionals. We also asked participants whether they thought they had received active treatment to examine whether the significant benefits seen with ethyl-EPA could be attributed to their guessing correctly their group allocation. We tried to approximate ordinary clinical practice by allowing treating physicians to make changes to participants’ medication when clinically required. Finally, we analysed the data on an intent-to-treat basis and showed a superior response to ethyl-EPA compared with placebo in spite of the difficulties in finding clear drug-placebo separation in add-on trials (Keck et al., 2000). This is particularly relevant here since about half of those randomised to the placebo group had their medication adjusted when their symptoms persisted or worsened.

### Table 1 Demographic and clinical characteristics of the 75 study participants

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Placebo (n = 26)</th>
<th>1 g/day ethyl-EPA (n = 24)</th>
<th>2 g/day ethyl-EPA (n = 25)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age, years: mean (s.d.)</strong></td>
<td>46.5 (10.3)</td>
<td>49.2 (11.7)</td>
<td>45.5 (9.6)</td>
</tr>
<tr>
<td><strong>Female: male, n</strong></td>
<td>16:10</td>
<td>19:5</td>
<td>22:3</td>
</tr>
<tr>
<td><strong>Diagnosis, n</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bipolar disorder I</td>
<td>24</td>
<td>19</td>
<td>22</td>
</tr>
<tr>
<td>Bipolar disorder II</td>
<td>2</td>
<td>5</td>
<td>3</td>
</tr>
<tr>
<td><strong>Duration of episode at study entry, months: mean (s.d.)</strong></td>
<td>5.6 (3.0)</td>
<td>6.0 (2.6)</td>
<td>5.2 (2.9)</td>
</tr>
<tr>
<td><strong>Age at onset of first depressive episode, years: mean (s.d.)</strong></td>
<td>23.6 (8.4)</td>
<td>24.2 (10.3)</td>
<td>26.1 (9.1)</td>
</tr>
<tr>
<td><strong>Age at onset of first manic episode, years: mean (s.d.)</strong></td>
<td>29.1 (9.4)</td>
<td>31.6 (12.9)</td>
<td>32.7 (9.4)</td>
</tr>
<tr>
<td><strong>Depressive episodes in the preceding 12 months, n: mean (s.d.)</strong></td>
<td>1.3 (1.2)</td>
<td>1.5 (1.5)</td>
<td>1.2 (1.1)</td>
</tr>
<tr>
<td><strong>Manic episodes in the preceding 12 months, n: mean (s.d.)</strong></td>
<td>0.4 (1.1)</td>
<td>0.1 (0.8)</td>
<td>0.1 (0.3)</td>
</tr>
<tr>
<td><strong>Hypomanic episodes in the preceding 12 months, n: mean (s.d.)</strong></td>
<td>0.5 (1.1)</td>
<td>0.5 (0.9)</td>
<td>0.1 (0.3)</td>
</tr>
<tr>
<td><strong>Mixed episodes in the preceding 12 months, n: mean (s.d.)</strong></td>
<td>0.2 (0.4)</td>
<td>0.3 (0.5)</td>
<td>0.02 (0.2)</td>
</tr>
<tr>
<td><strong>Hospital admissions in the preceding 12 months, n: mean (s.d.)</strong></td>
<td>0.3 (0.6)</td>
<td>0.1 (0.3)</td>
<td>0.2 (0.5)</td>
</tr>
<tr>
<td><strong>Lifetime hospital admissions, n: mean (s.d.)</strong></td>
<td>4.3 (5.3)</td>
<td>3.6 (2.9)</td>
<td>2.9 (2.6)</td>
</tr>
<tr>
<td><strong>Participants with a lifetime history of psychosis within episodes, n (%)</strong></td>
<td>21 (81)</td>
<td>15 (63)</td>
<td>17 (68)</td>
</tr>
</tbody>
</table>

**EPA, eicosapentaenoic acid.**

### Table 2 Participants’ concomitant medication at the time of study entry

<table>
<thead>
<tr>
<th>Concomitant medication</th>
<th>Placebo (n = 26)</th>
<th>1 g/day ethyl-EPA (n = 24)</th>
<th>2 g/day ethyl-EPA (n = 25)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lithium</td>
<td>9 (34.6)</td>
<td>15 (62.5)</td>
<td>10 (40.0)</td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>7 (26.9)</td>
<td>3 (12.5)</td>
<td>4 (16.0)</td>
</tr>
<tr>
<td>Sodium valproate</td>
<td>2 (7.6)</td>
<td>4 (16.6)</td>
<td>3 (12.0)</td>
</tr>
<tr>
<td>Antipsychotic</td>
<td>12 (46.1)</td>
<td>2 (8.3)</td>
<td>7 (28.0)</td>
</tr>
<tr>
<td>Antidepressant</td>
<td>7 (26.9)</td>
<td>12 (50.0)</td>
<td>12 (48.0)</td>
</tr>
<tr>
<td>Benzodiazepines</td>
<td>2 (7.6)</td>
<td>7 (29.1)</td>
<td>3 (12.0)</td>
</tr>
<tr>
<td>None</td>
<td>5 (19.2)</td>
<td>5 (20.8)</td>
<td>1 (4.0)</td>
</tr>
</tbody>
</table>

**EPA, eicosapentaenoic acid.**

### Table 3 Scores on the HRSD, YMRS and CGI at study entry and at end-point

<table>
<thead>
<tr>
<th>Scale</th>
<th>Placebo (n = 26)</th>
<th>1 g/day ethyl-EPA (n = 24)</th>
<th>2 g/day ethyl-EPA (n = 25)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>HRSD score: mean (s.d.)</strong></td>
<td>15.4 (5.0)</td>
<td>14.7 (4.3)</td>
<td>14.8 (5.6)</td>
</tr>
<tr>
<td><strong>YMRS score: mean (s.d.)</strong></td>
<td>6.3 (6.7)</td>
<td>6.7 (7.6)</td>
<td>6.6 (6.7)</td>
</tr>
<tr>
<td><strong>CGI score: mean (s.d.)</strong></td>
<td>3.0 (0.9)</td>
<td>3.0 (1.1)</td>
<td>2.9 (1.1)</td>
</tr>
</tbody>
</table>

**EPA, eicosapentaenoic acid; HRSD, Hamilton Rating Scale for Depression; YMRS, Young Mania Rating Scale; CGI, Clinical Global Impression Scale.**

### Possible mechanism of action of ethyl-EPA

The precise mechanism of action of ethyl-EPA in improving bipolar depression is not clear. Antidepressants exert their action at the level of neurotransmitters (catecholamines and serotonin) and neurotransmitter
The role of ethyl-EPA in bipolar disorder

This is the first randomised double-blind placebo-controlled clinical trial of ethyl-EPA in depression in people with bipolar disorder. Our results confirm initial observations (Horrobin & Peet, 2001; Nemets et al., 2002) of the antidepressant effect of omega-3 fatty acids, particularly of ethyl-EPA. They also strongly suggest that treatment with ethyl-EPA is not associated with increased risk of inducing manic symptoms. At the doses prescribed here the side-effects were minimal and indistinguishable from those in the placebo group. Although the role of ethyl-EPA in the treatment of bipolar disorder requires further evaluation, our results offer optimism that ethyl-EPA represents a new generation of naturally occurring and safe psychotropic compounds.

REFERENCES


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CLINICAL IMPLICATIONS

■ Adjunctive ethyl-eicosapentaenoic acid (EPA) treatment of bipolar depression is safe and well tolerated.

■ Adjunctive ethyl-EPA treatment appears to have antidepressant effects and minimal propensity to induce mania.

■ As ethyl-EPA is a naturally occurring compound it may prove more acceptable to patients than other pharmacological interventions.

LIMITATIONS

■ This small study only assessed short-term efficacy and tolerability of ethyl-EPA treatment in bipolar disorders; its value in long-term treatment is unknown.

■ This study only assessed the efficacy and tolerability of ethyl-EPA as adjunctive treatment in bipolar disorder; its value as monotherapy is unknown.

■ This study did not assess the efficacy of ethyl-EPA in severe bipolar depression.
Efficacy of ethyl-eicosapentaenoic acid in bipolar depression: randomised double-blind placebo-controlled study
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